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(S) Method of making highly adsorptive copolymers.

© A precipitation polymerization process for the production of a macroporous cross-linked copolymer in which there is added to a reactor equipped with a stirrer, at least one monounsaturated monomer and at least one polyunsaturated monomer, adding to the reactor a solvent in which the monomers are soluble but in which the copolymer is insoluble and conducting the copolymerization in the reactor by stirring the solvent, the monounsaturated monomer and the polyunsaturated monomer, at stirring rates in the reactor of from zero revolutions per minute to about three hundred revolutions per minute.

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METHOD OF MAKING HIGHLY ADSORPTIVE COPOLYMERS

In accordance with the present invention, copolymer powders are produced by novel processes not believed to be taught in the prior art. In general, the prior art relates to suspension polymerization processes for the production of porous polymeric and copolymeric spheres and beads in which the precipitant is present during polymerization. These are defined as an "in situ" process.

According to the prior art, crosslinked porous copolymers in particle form can be produced by at least three distinct processes. One process produces beads by "in situ" suspension polymerization. Another process produces beads by suspension polymerization but the beads are "post adsorbed" with an active ingredient after the volatile porogen is removed. In a third process, powders are produced by "in situ" precipitation polymerization.

What has been accomplished in accordance with the present invention, however, is a unique concept differing from all of the foregoing methods, and wherein post adsorbent powders are produced in a novel fashion.

This invention relates to a an improvement in the process for the production of a macroporous, cross-linked, copolymer in a reactor equipped with a stirrer, by precipitation polymerization in a solvent containing at least one monounsaturated monomer and at least one polyunsaturated monomer soluble therein and in which the improvement involves conducting the polymerization in the reactor at stirring rates of from zero revolutions per minute to about three hundred revolutions per minute.

The solvent can be removed from the porous copolymer at the conclusion of the polymerization, in which case, the copolymer is mixed with a functional material in order to disperse and uniformly distribute the functional material throughout the porous copolymer and to entrap the functional material therewithin. The functional material can be water, silicone oils, glycerine, mineral oil, organic esters, glycols and glutamates.

One monomer of the copolymer is a monounsaturated monomer such as lauryl methacrylate and the other monomer of the copolymer is a polyunsaturated monomer such as ethylene glycol dimethacrylate. The copolymer can also be formed using only polyunsaturated monomers. The copolymer is in the form of a powder and the powder is a combined system of particles. The system of powder particles includes unit particles of less than about one micron in average diameter, agglomerates of fused unit particles of sizes in the range of about twenty to eighty microns in average diameter and aggregates of clusters of fused agglomerates of sizes in the range of about two hundred to about twelve hundred microns in average diameter.

A precipitation polymerization process is used for producing the macroporous cross-linked copolymer. In the process, there is copolymerized at least one monounsaturated monomer and at least one polyunsaturated monomer in the presence of an organic liquid which is a solvent for the monomers but not for the copolymer. The process can also be conducted using only polyunsaturated monomers. The copolymerization of the monomers is initiated by means of a free radical generating catalytic compound, precipitating a copolymer in the solvent in the form of a powder. A dry powder is formed by removing the solvent from the precipitated copolymeric powder.

The solvent is preferably isopropyl alcohol, although ethanol, toluene, heptane, xylene, hexane, ethyl alcohol and cyclohexane, may also be employed. The monounsaturated monomer and the polyunsaturated monomer can be present in mol ratios of, for example, 20:80, 30:70, 40:60 or 50:50. The process includes the step of stirring the monomers, solvent and the free radical generating catalytic compound, during copolymerization. Preferably, the dry powder is formed by filtering excess solvent from the precipitated powder and the filtered powder is vacuum dried. The powder may then be "post adsorbed" with various functional materials.

The powders of the present invention may be used as carriers or adsorbents for materials such as water, aqueous systems, emollients, moisturizers, fragrances, dyes, pigments, flavors, drugs such as ibuprofen, phosphoric acid, insect repellents, vitamins, sunscreens, detergents, cosmetics, pesticides, pheromones, herbicides, steroids, sweeteners, pharmaceuticals and antimicrobial agents. Finely divided solids such as analgesic materials can be adsorbed by dissolving the finely divided analgesic in a solvent, mixing the analgesic and solvent with the powder and removing the solvent. Other post adsorbable materials include alkanes, alcohols, acid esters, silicones, glycols, organic acids, waxes and alcohol ethers.

These and other objects, features and advantages, of the present invention will become apparent when considered in light of the following detailed description, including the accompanying drawings.

Figure 1 is a photomicrograph of the various components of the complex structure of the powder produced in Example I and including unit particles, agglomeratures and aggregates.

Figures 2 and 3 are photomicrographs of the agglomerates and aggregates of Figure 1, respectively, shown on a larger scale.

Figure 4 is a photomicrograph of a polymer bead produced by suspension polymerization.

Figure 5 is a photomicrograph of the bead of Figure 4 with a portion of the shell removed to reveal the interior structure of the bead.

Figure 6 is a photomicrograph of a copolymeric powder material. The powder is shown in magnification as it appears when the agitation rate employed in the process for producing the powder is zero rpm.

Figures 7-10 are additional photomicrographs of copolymeric powder materials. The powder is shown in magnification as it appears when the agitation rate employed in the process for producing the powder varies from seventy-five rpm up to eight hundred rpm.

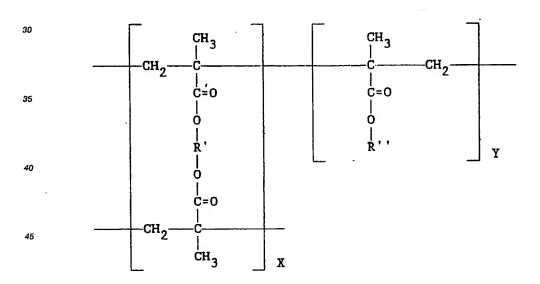
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In the above figures in the drawing, the magnification is indicated in each instance. For example, the magnification in Figures 6-9 is 1000X and 2000X in Figure 10. Figures 6-10 also include an insert identifying a length approximating ten microns for comparative purposes.

It should be pointed out, that in viewing the various figures, one will note that as the rate of stirring is increased from zero rpm up to eight hundred rpm, that the size of the unit particles increase. This is in direct opposition to what has been traditionally observed in suspension polymerization systems, wherein increases in stirring rates decrease particle size. Because of the increased size of the unit particles shown in Figure 10 and the resulting decrease in surface area, the adsorptive capacity of these large particles is less than the adsorptive capacity of the smaller sized particles shown in Figures 6-9.

The most effective unit particles can be produced if the rate of stirring is maintained below about three hundred rpm, although particles produced at rates beyond three hundred rpm are useful and adsorptive, but to a lesser extent.

The material of the present invention, can be broadly and generally described as a crosslinked copolymer capable of entrapping solids, liquids and gases. The copolymer is in particulate form and constitutes free flowing discrete solid particles even when loaded with an active material. When loaded, it may contain a predetermined quantity of the active material. One copolymer of the invention has the structural formula:



where the ratio of x to y is 80:20, R' is -CH₂CH₂- and R" is -(CH₂)₁₁CH₃.

The copolymer is a highly cross linked copolymer, as evidenced by the foregoing structural formula and is more particularly a highly crosslinked polymethacrylate copolymer. This material is manufactured by the Dow Corning Corporation, Midland, Michigan, U.S.A., and sold under the trademark POLYTRAP®. It is a low density, highly porous, free-flowing white particulate and the particles are capable of adsorbing high levels of lipophilic liquids and some hydrophilic liquids, while at the same time maintaining a free-flowing particulate character.

In the powder form, the structure of the particulate is complex and consists of unit particles less than one micron in average diameter. The unit particles are fused into agglomerates of twenty to eighty microns

in average diameter. Thes agglomerates are loosely clustered into macro-particles termed aggregates of about 200 to about 1200 microns in average diameter.

Adsorption of actives to form post adsorbent powder, can be accomplished using a stainless steel mixing bowl and a spoon, wherein the active ingredient is added to the empty dry powder and the spoon is used to gently fold the active into the powder. Low viscosity fluids may be adsorbed by addition of the fluids to a sealable vessel containing the powder and tumbling the materials until a consistency is achieved. More elaborate blending equipment such as ribbon or twin cone blenders can also be employed.

The following example illustrates the method for making a post adsorbent powder, of the type illustrated in Figures 1-3 and 6-10.

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Example I

.A hydrophobic porous copolymer was produced by the precipitation polymerization technique by mixing in a five hundred milliliter polymerization reactor equipped with a paddle type stirrer, 13.63 grams of ethylene glycol dimethacrylate monomer, or eighty mole percent, and 4.37 grams of lauryl methacrylate monomer, or twenty mole percent. Isopropyl alcohol was added to the reactor as the solvent in the amount of 282 grams. The monomers were soluble in the solvent, but not the precipitated copolymer. The process can be conducted with only polyunsaturated monomers if desired. The mixture including monomers, solvent and 0.36 grams of catalytic initiator benzoyl peroxide, was purged with nitrogen. The system was heated by a water bath to about 60°C. until copolymerization was initiated, at which time, the temperature was increased to about 70-75° C. for six hours, in order to complete the copolymerization. During this time, the copolymer precipitated from the solution. The copolymerization produced unit particles of a diameter less than about one micron. Some of the unit particles adhered together providing agglomerates of the order of magnitude of about twenty to eighty microns in diameter. Some of the agglomerates adhered further and were fused and welded one to another, forming aggregates of loosely held assemblies of agglomerates of the order of magnitude of about two to eight hundred microns in diameter. The mixture was filtered to remove excess solvent and a wet powder cake was tray dried in a vacuum oven. A dry hydrophobic copolymeric powder consisting of unit particles, agglomerates and aggregates was isolated.

The adsorptive capacity of the hydrophobic particulates produced in Example I, as a function of the stirring rate, was determined. The stirring rate during the reaction in Example I significantly influenced the adsorption properties of the particulate materials. The adsorptivity of the particulate materials decreases with an increase in stirring rate and the density of the particulates increases. These results are set forth in Tables I-III.

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TABLE !

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Agitation Rate (RPM)	Bulk Density Size (g/cc)	Average Aggregate Size (µ)	Average Agglomerate Size (µ)	Average Unit Particle Size (μ)	Adsorption Capacity*
0 75 150 300 800	0.067 0.077 0.071 0.293 0.440	182.5 140.6 149.8 47.0	33.9 36.6 39.8 34.0 10.0	1.0 0.5 0.8 1.5-2.0 3.0-5.0	83.0 84.8 83.0 58.3 37.7

^{* =} Percent Silicone Oil

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TABLE II

	Adsorption Capacity %						
Stirring Speed RPM	Water	Mineral Oil	Glycerine	Organic Ester*			
0 75 150 300	0 0 0	80 83.9 80 54.5	75 75 75 58.3	80 81.5 80 54.5			

^{* = 2-}ethylhexyl-oxystearate

RPM

1000

Mineral

Oil

32.8

15

10

TABLE III

Silicone

Οii

32.9

Density (g/cm3)

Tapped

0.5261

Bulk

0.3808

20

25

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0.0580 n 82.5 82.5 86.5 0.0368 82.2 75 82.3 86.5 0.0462 0.0667 150 82.3 82.3 86.3 0.0527 0.0737 200 81.5 81.5 85.7 0.0554 0.0752 0.0636 0.0859 250 79.2 0.08 84.8 300 68.8 68.8 75.0 0.1300 0.1768 58.3 61.5 0.1736 0.2392 450 58.3 600 54.5 54.5 60 0.1933 0.2792 700 42.2 42.5 45.7 0.2778 0.4142 800 33.3 28.6 33.3 0.3862 0.5322

Adsorption Capacity %

2-ethylhexyl

oxystearate

28.5

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In the foregoing tables, it can be seen that adsorption and density, as a function of stirring rate, was determined for several fluids including a silicone oil, water, mineral oil, glycerine and an organic ester. From zero rpm up to about 250 rpm, the adsorptivity of the porous copolymeric powder particulates of Example I remained essentially consistent. However, at about three hundred rpm, there was a substantial decrease in adsorptivity, which decrease became more apparent as the stirring rate was increased up to about one thousand rpm. A similar pattern is evidenced by the data which are reflective of the density.

This phenomenon is more apparent in the photomicrographic figures of the drawing. Thus, it can be seen from Figure 6, that the particle size of the unit particles increases as the stirring rate is increased, as evidenced by Figure 10. A progression in this phenomenon can be observed in Figures 7-9.

While the procedure of Example I is a precipitation polymerization process and not a suspension polymerization system, the prior art dealing with suspension polymerization processes, teaches that an increase in stirring rate causes a decrease in particle size. This is documented, for example, in U.S. Patent No. 4,224,415, issued September 23, 1980, and in the PCT International Publication No. WO 88/01164, published on February 25, 1988. The PCT International Publication employs stirring rates upwards of nine hundred to twelve hundred rpm. In Example I of the present invention, however, increases in stirring rates not only did not decrease the particle size, but in fact had exactly the opposite effect, causing the unit particle size to increase. As the rate of stirring increased from zero rpm up to one thousand, the density of the particles increased and the adsorptive capacity decreased.

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In accordance with the above, it is possible to tailor porous adsorbent powders of a particular particle size and adsorptivity by means of stirring rate. Thus, with large unit particles in Figure 10, the adsorptive capacity is less than the adsorptive capacity of smaller sized unit particles in Figures 6-9. While the most effective particles are produced when the rate of stirring is maintained below about three hundred rpm, particles produced at rates beyond three hundred rpm are useful.

It is important to understand that the method of Example I for the production of porous copolymer particulate powder materials is characterized as a precipitation polymerization technique. In accordance with the technique, monomers are dissolved in a compatible volatil solvent in which both monomers are soluble. Polymer in the form of a powder is precipitated and the polymer is insoluble in the solvent. No surfactant or dispersing aid is required. The materials produced are powders and not spheres or beads. The powder particulates include unit particles, agglomerates and aggregates. The volatile solvent is subsequently removed resulting in a dry powder, which can be post adsorbed with a variety of functional active ingredients. The suspension polymerization process on the other hand, provides that polymerization be carried out in water and in some cases chloroform or chlorinated solvents. The monomers, the active and the catalyst, form beads or droplets in water and polymerization occurs within each bead. A surfactant or stabilizer, such as polyvinyl pyrrolidone, is required in order to prevent the individually formed beads and droplets from coalescing. The resulting beads, with the active material entrapped therein, include a substantially spherical outer crust or shell, the interior of which contains a macroporous structure of fused unit particles, agglomerates and aggregates. The bead is about ten microns in average diameter to about one hundred-fifty microns, depending upon the rate of agitation employed during the process. Such beads are shown in Figures 4 and 5.

Some unique features of the powders of Example I and Figures 1-3 and 6-10 are their ability to adsorb from sixty to eighty percent of a liquid and yet remain free flowing. The materials provide a regulated release of volatile ingredients such as cyclomethicone entrapped therein and have the capability of functioning as carriers for other non-volatile olls. Loaded powders disappear when rubbed upon a surface. This phenomenon is believed due to the fact that large aggregates of the material scatter light rendering the appearance of a white particulate, however, upon rubbing, these large aggregates decrease in size approaching the range of visible light and hence seem to disappear. The materials find applications in diverse areas such as cosmetics and toiletries, household and industrial products, pesticides, pheromone carriers and pharmaceuticals. The materials do not swell in common solvents and are capable of physically adsorbing active ingredients by the filling of interstitial voids by capillary action. The active ingredients are subsequently released by capillary action or wicking from the voids within the particulates.

The following example illustrates a precipitation polymerization process in which an organic ester is entrapped "in situ" in the polymer powder.

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Example II

7 grams of 2-ethylhexyl oxystearate was mixed with 1.5 grams of ethylene glycol dimethacrylate and 1.5 grams of lauryl methacrylate in a glass test tube. The solution was deaerated for five (5) minutes and 0.1 ml of t-butyl peroctoate was added and mixed while heating to 80°C. in an oil bath. After 20 minutes, the contents solidified; and the mixture was maintained at about 80°C. for an additional hour to assure full polymerization. A semi-soft, heterogeneous white opaque polymer mass resulted containing the entrapped ester.

The powder of Example II differs from the powder of Example I in that the solvent in Example I is removed resulting in a dry empty powder which is post adsorbed with other functional materials. The powder of Example II is otherwise similar to the material shown in Figures 1-3.

Example III illustrates a process for the production of beads as shown in Figures 4 and 5. The process is suspension polymerization and an organic ester is entrapped "in situ".

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Example III

1.20 grams of polyvinyl pyrrolidone was dissolved in 1500 ml of water in a 2000 ml three necked resin flask equipped with a stirrer, thermometer and nitrogen purge. A solution of 335 grams of 2-ethylhexyl oxystearate, 132 grams ethylene glycol dimethacrylate, 33 grams 2-ethylhexyl methacrylate and 5 ml thutyl peroctoate, was bubbled with nitrogen for 5 minutes. The resultant mix was slowly added to the stirred aqueous solution of polyvinyl pyrrolidone at 22° C. under nitrogen. The temperature was raised to 80° C. with constant agitation and held until polymerization started in approximately 15 minutes and maintained at 80° C. for an additional 2 hours to complete the reaction. Semi-soft, white opaque beads were collected by filtering off the supernatant liquid and dried to remove any excess water. The beads weighed 450 g for a yield of 90% and were 0.25 to 0.5 mm in diameter. Other protective colloids such as starch, polyvinyl alcohol, carboxymethyl cellulose, methyl cellulose or inorganic systems such as divalent alkali metal

hydroxides, for xample, MgOH may be used in place of the polyvinyl pyrrolidon suspending medium.

In Example III macroporous polymers submicron in size are produced with two or more monomers, at least one monomer of which contains more than a single double bond. The polymerization is conducted in the presence of an active ingredient which does not dissolve or swell the resulting polymer. The monomers and the active ingredient are mutually soluble, but are insoluble in the aqueous suspending medium in which droplets are formed. Polymerization occurs within suspended droplets and beads or spheres are produced. The active ingredient which is polymerized "in situ" is entrapped and contained within the beads, but the active ingredient is capable of being released. It is also possible to use a volatile liquid during polymerization and to subsequently thermally drive off the volatile liquid, leaving behind a porous polymer bead product into which a variety of active materials can be subsequently adsorbed.

Examples of polyunsaturated monomers suitable for use in accordance with the present invention are ethylene glycol dimethacrylate, triethylene glycol dimethacrylate, tetraethylene glycol dimethacrylate, trimethylol propane ethoxylated triacrylate, ditrimethylol propane dimethacrylate; propylene, dipropylene and higher propylene glycols, 1,3-butylene glycol dimethacrylate, 1,4 butanediol dimethacrylate, neopentyl glycol dimethacrylate, pentaerythritol dimethacrylate, dipentaerythritol dimethacrylate, bisphenol A dimethacrylate, divinyl and trivinyl benzene, divinyl and trivinyl toluene triallyl maleate, triallyl phosphate, diallyl maleate, diallyl itaconate and allyl methacrylate. The mono-unsaturated monomers include allyl methacrylates and acrylates having straight or branched chain alkyl groups with 1 to 30 carbon atoms, preferably 5 to 18 carbon atoms. Preferred monomers include lauryl methacrylate, 2-ethylhexyl methacrylate, isodecylmethacrylate, stearyl methacrylate, hydroxy ethyl methacrylate, hydroxy propyl methacrylate, diacetone acrylamide, phenoxy ethyl methacrylate, tetrahydrofurfuryl methacrylate and methoxy ethyl methacrylate.

As noted previously, the copolymer can be formed by copolymerizing one monounsaturated monomer with one polyunsaturated monomer or with only polyunsaturated monomers.

Example IV

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Example I was repeated for each of a series of monomer systems shown in Tables IV-XVII. In each instance, submicron sized copolymeric powders were produced employing a stirring speed of about seventy-five RPM. The catalyst was dibenzoyl peroxide. Adsorption capacities of the various copolymeric powders for fluids were determined and are shown in the Tables, along with the mole ratios of monomers and the solvent. The abbreviations used in Tables IV-XVII are identified as follows:

DAA Diacetone acrylamide

EGDM Ethylene gylcol dimethacrylate
TEGDM Tetraethylene glycol dimethacrylate
ST Styrene
DVB Divinylbenzene
VP Vinyl pyrrolidone

O IBOMA Isobornyl methacrylate
PEMA Phenoxyethyl methacrylate
IDMA Isodecyl methacrylate
STMA Stearyl methacrylate
HPMA Hydroxypropyl methacrylate

CYMA Cyclohexyl methacrylate
DMAEMA Dimethylaminoethyl methacrylate
TBAEMA t-butyl aminoethyl methacrylate
AMPS 2-acrylamido propane sulfonic acid
BMA Butyl methacrylate

EHMA 2-ethylhexyl methacrylate MMA Methyl methacrylate HEMA 2-hydroxyethyl methacrylate EHO 2-ethylhexyl oxystearate GG Glucose glutamate

FEG Polyethylene glycol 200

TABLE IV

Adsorption Capacity % Monomers Mole Solvent **EHO** Glycerine GG Water Ratio DAA/EGDM 20/80 Xylene 83 78 DAA/EGDM 30/70 Xylene 77 80 83 78 DAA/EGDM 40/60 Xylene 75 75 83 77 DAA/EGDM 50/50 **Xyiene** 50 57 67 0 **DAA/EGDM** 60/40 Xylene 40 40 50 0 Xylene DAA/TEGDM 20/80 40 50 62 58 DAA/TEGDM 30/70 Xylene 29 40 50 55 DAA/TEGDM 40/60 Xylene 25 28 40 43 DAA/TEGDM 50/50 **Xylene** 25 30 40 43 DAA/TEGDM 60/40 Xylene 22 29 40 40

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TABLE V

Adsorption Capacity % 25 Monomers Mole Solvent EHO Glycerine PEG Water Ratio ST/TEGDM 20/80 **IPA** 69 69 67 ST/TEGDM 30/70 **IPA** 58 64 67 69 ST/TEGDM 40/60 **IPA** 62 71 71 61 30 ST/TEGDM 50/50 IPA 67 62 54 58 ST/TEGDM 60/40 **IPA** 50 58 58 54 ST/TEGDM 70/30 **IPA** 50 58 50 54 ST/TEGDM 80/20 **IPA** 44 54 50 50 ST/DVB 20/80 **IPA** 80 .75 75 0 35 ST/DVB 30/70 **IPA** 75 67 75 0 ST/DVB 40/60 **IPA** 69 67 67 0 ST/DVB 50/50 **IPA** 64 72 67 0 ST/DVB 60/40 **IPA** 67 71 71 0 ST/DVB 70/30 **IPA** 71 75 76 0 40

IPA

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50

50

0

80/20

45

ST/DVB

50

TABLE VI

	[Adsorption Capacity %				
Monomers	Mole Ratio	Solvent	EHO	Glycerine	GG	Water	
VP/EGDM	20/80	Xylene	77	80	7.4	73.6	
VP/EGDM	30/70	Xylene	76	79	78.3	70	
VP/EGDM	40/60	Xylene	70	67	75.6	75	
VP/EGDM	50/50	Xylene	72	76	80	76	
VP/EGDM	60/40	Xylene	74	80	76	77	
VP/EGDM	70/30	IPA	71	78	74	75	
VP/EGDM	80/20	IPA	67	75	73	74	
VP/TEGDM	20/80	Xylene	58	68.8	61.5	67.7	
VP/TEGDM	30/70	Xylene	70	67	54.5	68.8	
VP/TEGDM	40/60	Xylene	54.5	61.5	52.4	64.3	
VP/TEGDM	50/50	Xylene	44.4	47.4	52.4	52.4	
VP/TEGDM	60/40	Xylene	50	44.4	50	54.4	
VP/TEGDM	70/30	Xylene	50	47.4	44.4	50	
VP/TEGDM	80/20	Xylene	54.5	52.4	60	58	

TABLE VII

						%
Monomers	Mole Ratio	Solvent	EHO	Glycerine	GG	Water
IBOMA/EGDM	20/80	IPA	62.9	9.1	71.4	0
IBOMA/EGDM	30/70	IPA	64.3	16.6	67.7	0
IBOMA/EGDM	40/60	IPA	68.7	28.6	61.5	0
IBOMA/EGDM	50/50	IPA	67.7	16.7	58.3	0
IBOMA/EGDM	60/40	IPA	50	23.1	50	0
IBOMA/EGDM	70/30	IPA	50	9.1	47.3	0
IBOMA/EGDM	80/20	IPA	52.3	16.6	44.4	0
IBOMA/TEGDM	20/80	IPA	66.6	62.9	61.5	0
IBOMA/TEGDM	30/70	IPA	61.5	61.5	70.6	0
IBOMA/TEGDM	40/60	IPA	64.3	64.3	71.4	0
IBOMA/TEGDM	50/50	IPA	61.5	66.6	67.7	0
IBOMA/TEGDM	60/40	IPA	58.3	54.5	54.5	0
IBOMA/TEGDM	70/30	IPA	47.3	50	41.1	0
IBOMA/TEGDM	80/20	IPA	37.5	41.1	33.3	0

TABLE VIII

Adsorption Capacity % Mole Solvent Monomers **EHO** Glycerine Water Ratio PEMA/EGDM 20/80 IPA 64.3 68.7 66.6 61.5 PEMA/EGDM 30/70 **IPA** 54.5 50 54.5 44.4 PEMA/EGDM 40/60 **IPA** 52.3 47.3 72.2 9 PEMA/EGDM 50/50 **IPA** 54.5 33.3 62.9 0 PEMA/EGDM 60/40 **IPA** 67.7 28.5 70.5 0 **PEMA/EGDM** 70/30 IPA · 69.7 44.4 60.7 0 **PEMA/EGDM** 80/20 **IPA** 66.6 68.7 66.6 0 PEMA/TEGDM 20/80 **IPA** 58.3 56.5 66.6 58.3 PEMA/TEGDM 30/70 **IPA** 64.2 70.5 67.7 62.9 PEMA/TEGDM 40/60 **IPA** 66.6 67.7 71.4 69.7 PEMA/TEGDM 50/50 **IPA** 66.6 70.5 73.6 72.2 **PEMA/TEGDM** 60/40 **IPA** 58.3 62.9 52.3 61.5 PEMA/TEGDM 70/30 **IPA** 58.3 50 52.3 54.5 PEMA/TEGDM 80/20 **IPA** 67.7 73.6 76.1 47.3

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TABLE IX

			Adsorption Capacity %				
Monomers	Mole Solvent Ratio		EHO	Glycerine	PEG	Water	
IDMA/EGDM	20/80	IPA	55	64	70	0	
IDMA/EGDM	30/70	IPA	38	50	44	0	
IDMA/EGDM	40/60	. IPA	50	67	69	0	
IDMA/EGDM	50/50	IPA	58	64	67	0	
IDMA/EGDM	60/40	IPA	58 ·	69	69	0	
IDMATEGDM	20/80	IPA	62	70	70	0	
IDMA/TEGDM	30/70	IPA	50	62	62	0	
IDMA/TEGDM	40/60	IPA	62	67	67	0	
IDMA/TEGDM	50/50	IPA	38	44	50	0	
IDMA/TEGDM	60/40	IPA	38	55	50	0	

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TABLE X

Adsorption Capacity % Monomers Mole Solvent EHO Water Glycerine PEG Ratio STMA/EGDM 10/90 **IPA** 66 64.3 66.7 0 STMA/EGDM 20/80 **IPA** 69 63 65.5 0 **STMA/EGDM** 30/70 **IPA** 73-75 58.3 61.5 0 STMA/EGDM 40/60 **IPA** 69-71 54.5 58.3 0 STMA/EGDM 50/50 **IPA** 60-63 52.4 52.4 0 STMA/TEGDM 20/80 **IPA** 50 47.4 52.4 0 **STMA/TEGDM** 30/70 **IPA** 50 64.3 50 0 STMA/TEGDM 40/60 IPA 52.4 61.5 58.3 0 52.4 STMA/TEGDM **IPA** 50/50 47.4 56.5 0

TABLE XI

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Adsorption Capacity % Monomers Mole Solvent EHO Glycerine PEG Water Ratio **HPMA/EGDM** 20/80 **Xylene** 64.3 61.5 61.5 9 HPMA/EGDM 30/70 Xylene 54.5 16.7 58.3 0 HPMA/EGDM 40/60 Xylene 54.5 58.3 0 HPMA/EGDM 50/50 Xylene 37.5 58.3 50 0 HPMA/EGDM 60/40 Xylene 44.4 61.5 58.3 0 HPMA/EGDM 70/30 Xylene 37.5 50 44.4 0 **HPMA/EGDM** 80/20 Xylene 61.5 16.7 58.3 0 HPMA/TEGDM 20/80 Xylene 54.5 50 58.3 61.5 **HPMA/TEGDM** 30/70 Xylene 56.5 54.5 50 60 HPMA/TEGDM 40/60 Xylene 50 58.3 52.4 54.5 HPMA/TEGDM 50/50 **Xylene** 52.4 61.5 · 54.5 56.5 HPMA/TEGDM 60/40 Xylene 33.3 47.4 44.4 54.5 HPMA/TEGDM 70/30 Xylene 54.5 44.4 54.5 50 **HPMA/TEGDM** 80/20 **Xylene** 50 47.4 41.2 37.5

TABLE XII

Adsorption Capacity % Water PEG **EHO** Glycerine Monomers Mole Solvent Ratio **IPA** 66.6 CYMA/EGDM 80/20 61.5 71.4 0 0 70/30 **IPA** 66 64.2 CYMA/EGDM 60 66.6 0 60/40 **IPA** 66 CYMA/EGDM 61.5 CYMA/EGDM 50/50 **IPA** 66 68.7 0 64.2 68.7 0 CYMA/EGDM 40/60 **IPA** 64.2 66 0 66 66.6 **IPA** CYMA/EGDM 30/70 61.5 61.5 66.6 71.4 20/80 **IPA** 75 CYMA/EGDM **IPA** 68.7 0 68.7 0 80/20 CYMA/TEGDM 0 0 CYMA/TEGDM 70/30 **IPA** 71.4 69.7 60/40 0 62.9 0 **CYMA/TEGDM IPA** 66.6 50/50 0 0 CYMA/TEGDM **IPA** CYMA/TEGDM 40/60 **IPA** 60 0 72.9 0 CYMA/TEGDM 30/70 **IPA** 64.2 0 72.2 0 **CYMATEGDM** 20/80 **IPA** 61.5 0 66.6 0

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TABLE XIII

Adsorption Capacity % Glycerine **EHO** Mole Solvent Water Mineral Monomers Ratio Oil DMAEMA/EGDM 20/80 58.3 66.7 58.3 Hexane 0 DMAEMA/EGDM 40/60 Hexane 66.7 61.5 70.6 66.7 DMAEMA/EGDM 60/40 Hexane 77.3 61.5 72.2 76.2 68.8 58.3 DMAEMA/EGDM 80/20 Hexane 66.7 58.3 70.6 75 70.6 TBAEMA/EGDM 20/80 Hexane 0 TBAEMA/EGDM 40/60 Hexane 0 66.7 72.2 66.7 68.75 61.5 TBAEMA/EGDM 60/40 Hexane 0 61.5 54.6 50 80/20 44.4 TBAEMA/EGDM Hexane 0 80/20 54.6 54.6 58.3 50 Hexane TBAEMA/EGDM

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TABLE XIV

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			Adsorption Capacity %				
Monomers	Mole Solvent Ratio	Water	Mineral Oil	Glycerine	EHO		
AMPS/EGDM	20/80	Xylene	84.3	83.3	85.3	83.3	
BMA/EGDM	20/80	Hexane	0	70.6	75	68.8	
BMA/EGDM	40/60	Hexane	0	70.6	77.3	70.6	
BMA/EGDM	40/60	Ethyl-Alcohol	0	66.7	73.7	68.8	
BMA/EGDM	60/40	Hexane	0	72.2	0	73.7	
BMA/EGDM	80/20	Hexane	0	54.5	66.7	58.3	

TABLE XV

Adsorption Capacity % Water **EHO** Monomers Mole Solvent Mineral Glycerine Ratio Oil 2 EHMA/EGDM 20/80 **IPA** 68.8 66.7 64.3 0 2 EHMA/EGDM 30/70 **IPA** 0 68.8 66.7 64.3 70.6 2 EHMA/EGDM 40/60 **IPA** 66.7 66.7 0 2 EHMA/EGDM 50/50 **IPA** 64.3 68.3 61.5 0 2 EHMA/EGDM 60/40 **IPA** 61.5 64.3 50 0 2 EHMA/EGDM 70/30 **IPA** 0 58.3 64.3 50 2 EHMA/EGDM 80/20 **IPA** 58.3 64.3 50

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TABLE XVI

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			Adsorption Capacity %				
	Mole Solvent Ratio	Water	Mineral Oil	Glycerine	EHO		
MMA/EGDM MMA/EGDM MMA/EGDM MMA/EGDM MMA/EGDM MMA/EGDM MMA/EGDM	20/80 30/70 40/60 50/50 60/40 70/30 80/20	IPA IPA IPA IPA IPA IPA	61.5 64.3 61.5 58.3 54.5 66.7 66.7	58.3 61.5 64.3 54.5 50 61.5 44.4	64.3 66.7 64.3 61.5 61.5 72.2	58.3 61.5 61.5 58.3 54.5 64.3	

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TABLE XVII

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		Adsorption Capacity 9					
Monomers	Mole Ratio	Solvent	Water	PEG	Glycerine	EHO	
HEMA/EGDM HEMA/EGDM	20/80 30/70	IPA IPA	54.5 58.3	54.5 64.3	54.5 58.3	50 54.5	
HEMA/EGDM	40/60	IPA	58.3	61.5	64.3	54.5 54.5	
HEMA/EGDM HEMA/EGDM	50/50 60/40	IPA IPA	61.5 61.5	61.5 64.3	61.5 61.5	50 50	
HEMA/EGDM	70/30	IPA	58.3	64.3	58.3	50	
HEMA/EGDM	80/20	IPA	61.5	58.3	61.5	54.5	

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The water adsorbing porous polymeric materials produced above in some instances are to be contrasted with the water containing beads of U.S. Patent No. 3,627,708, issued December 14, 1971. The bead of the '708 patent is produced by "in situ" suspension polymerization and is adapted to contain water only because of the presence of a solubilizer such as sodium bis(2-ethyl hexyl) sulfosuccinate. The materials of Example IV, on the other hand, are produced by a precipitation polymerization process, which contains no solubilizer and produces a material in the form of a powder consisting of unit particles, agglomerates and aggregates. Thus, these materials are very distinct from the materials of the '708 patent.

It will be apparent from the foregoing that many other variations and modifications may be made in the

structures, compounds, compositions and methods described herein without departing substantially from the essential features and concepts of the present invention. Accordingly, it should be clearly understood that the forms of the invention described herein are exemplary only and are not intended as limitations of the scope of the present invention.

Claims

- 1. An improvement in the process for the production of a macroporous cross-linked copolymer in a reactor equipped with a stirrer by precipitation polymerization in a solvent containing at least one polyunsaturated monomer soluble therein, which improvement comprises conducting the polymerization in the reactor at stirring rates of from zero revolutions per minute.
- The process of claim 1 wherein the polyunsaturated monomer is selected from the group consisting
 of divinylbenzene, tetraethylene glycol dimethacrylate and ethylene glycol dimethacrylate.
 - 3. The process of claim 2 wherein the solvent is selected from the group consisting of isopropyl alcohol, xylene, hexane and ethyl alcohol.
 - The process of claim 1 wherein the solvent is removed from the copolymer at the conclusion of the polymerization.
 - 5. The process of claim 4 wherein the copolymer is mixed with a functional material in order to disperse and uniformly distribute the functional material throughout the copolymer and to entrap the functional material therewithin.
 - 6. The process of claim 5 wherein the functional material is selected from the group consisting of water, glycols, glutamates, silicone oils, glycerine, mineral oil and organic esters
 - 7. A precipitation polymerization process for the production of a macroporous cross-linked copolymer comprising adding to a reactor equipped with a stirrer at least one polyunsaturated monomer, adding to the reactor a solvent in which the monomer is soluble but in which the copolymer is insoluble and conducting the copolymerization in the reactor by stirring the solvent and the polyunsaturated monomer at stirring rates in the reactor of from zero revolutions per minute to about three hundred revolutions per minute.
 - 8. A precipitation polymerization process for the production of a macroporous cross-linked hydrophobic copolymer comprising adding to a reactor equipped with a stirrer at least one monounsaturated monomer and at least one polyunsaturated monomer, adding to the reactor a solvent in which the monomers are soluble but in which the copolymer is insoluble and conducting the copolymerization in the reactor by stirring the solvent, the monounsaturated monomer and the polyunsaturated monomer, at stirring rates in the reactor of from zero revolutions per minute to about three hundred revolutions per minute; the polyunsaturated monomer being selected from the group consisting of divinylbenzene, ethylene glycol dimethacrylate and tetraethylene glycol dimethacrylate; the monounsaturated monomer being selected from the group consisting of diacetone acrylamide, styrene, vinyl pyrrolidone, isobornyl methacrylate, phenoxyethyl methacrylate, isodecyl methacrylate, stearyl methacrylate, hydroxypropyl methacrylate, cyclohexyl methacrylate, dimethylaminoethyl methacrylate, t-butyl aminoethyl methacrylate, 2-acrylamido propane sulfonic acid, butyl methacrylate, 2-ethylhexyl methacrylate, methyl methacrylate, lauryl methacrylate and 2-hydroxyethyl methacrylate.

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FIG.I

1500 X

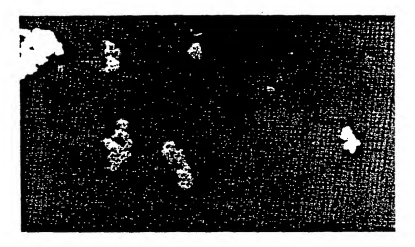


FIG.2

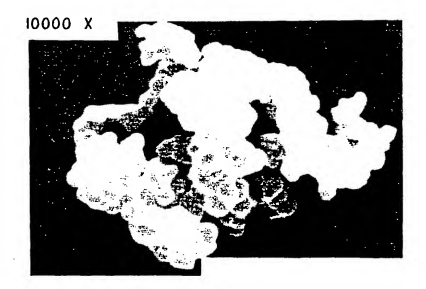


FIG.3

2000 X

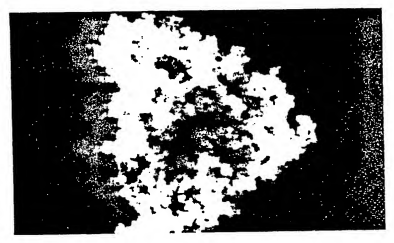


FIG.4

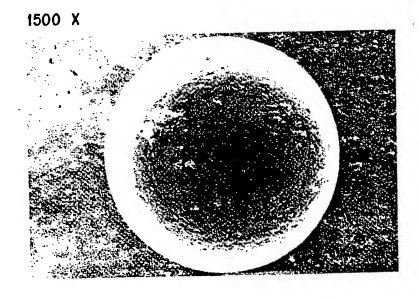
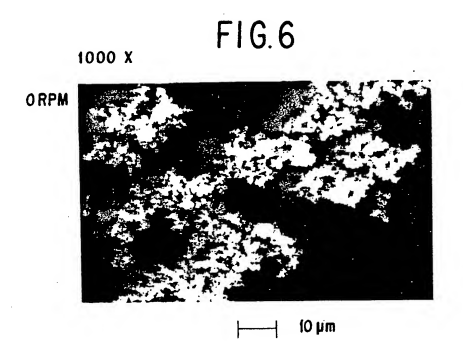


FIG. 5





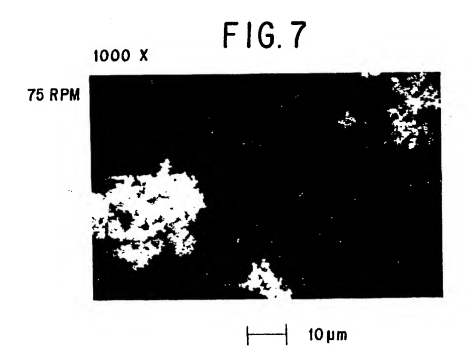


FIG.8

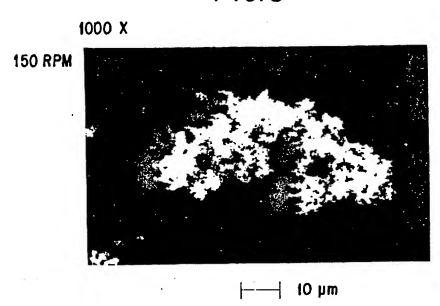


FIG.9

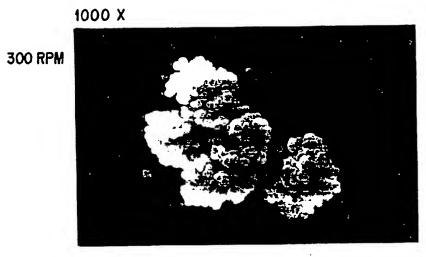
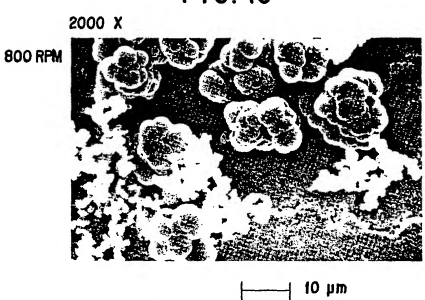


FIG. 10





EUROPEAN SEARCH REPORT

<u>.</u>	DOCUMENTS CONS	IDERED TO BE R	ELEVANT		EP	901	07026.8
Category		h indication where appropriant passages		elevant o claim			ION OF THE
х	40; page 16, line	4 021 5,6; page 11 12, line 2; 1 10 - page 18 lines 26-29	, line page , line 5;	-8	0000	08 F	220/00 212/00 226/10 2/44
A			mples;	-8			
A	GB - A - 2 146 (NIPPON SHOKUB KOGYO CO. LTD. * Claim 1; 40-47 *	AI KAGAKU	8	,2,7,			
D,A	US - A - 4 224 (MEITZNER et a		1 8	-3,7,			
	3; example 1	*				L FIELDS O (Int C ¹⁴)	
					0000	08 F	8/00 212/00 220/00 226/00
	The present search report has t	been drawn up for all claims					
 	Place of search	Date of completion		1	E	xaminer	
	VIENNA	23-07-199	0	P	USTE	RER	
Y : part doc A : tecl O : non	CATEGORY OF CITED DOCI ticularly relevant if taken alone ticularly relevant if combined w tument of the same category hnological background i-written disclosure trimediate document	rith another D L	: theory or princ earlier patent d after the filling document cited document cited member of the document	ocument, date d in the ap d for other	but pul plicatio reason	olished i in is	on, or